(11)

(12)

# FUROPEAN PATENT SPECIFICATION

- (45) Date of publication and mention of the grant of the patent 27.05.1998 Bulletin 1998/22
- (21) Application number, 95931175.4
- (22) Date of filing: 19.08,1995

- (51) Int CL6: C11D 11/00, C11D 1/02. C11D 1/14
- (86) International application number. PCT/FP95/03321
- (87) International publication number. WO 96/06916 (07.03.1996 Gazette 1996/11)
- (54) PRODUCTION OF ANIONIC SURFACTANT GRANULES

HESTELLUNG VON ANIONTENSIDGRANULATEN PRODUCTION DE GRANULES DE TENSIOACTIE ANIONIQUE

- (72) Inventors: (84) Designated Contracting States:
- (30) Priority: 26.08.1994 GB 9417356

BE DE ES ER GRIT

- (43) Date of publication of application: 11.06.1997 Bulletin 1997/24
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  - WO-A-93/19155

FP-A- 0 572 957

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#### Description

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The present invention relates to detergent particles, a process for their production and a composition containing them. More particularly the present invention relates to a process for the production of detergent particles having a high level of anionic surfactant which involves drying a detergent paste containing the anionic surfactant, and to the particles thereby obtained.

Detergent-active compounds conventionally employed in detergent compositions include anionic surfactants e.g. Intera railytipherance sulphonatos (LAS), linear alkyl other sulphate (LES) and primary alkyl sulphates (PAS); and nonionic surfactants e.g. alcohol ethoxylates. To improve detergency performance it is desirable to provide a high level of detergent-active material in the powder.

Often, the maximum level of active that may be incorporated is limited by process requirements. Detergent composition having a high bulk density are typically prepared by a process involving mixing or granulation of components of the composition and/or a base powder obtained for example from a spray-drying process and provide significant consumer benefits as compared to compositions of lower bulk density. It is known to incorporate detergent active compounds into such compositions in liquid from However as it is in excessing to control the retire of liquids to solid in order to form detargent granules the maximum level of detergent active material which may be incorporated in this manner is limited. It is also known to incorporate anionic surfactant a g. PAS in detargent compositions by means of a solid adjunct, that is, a particle comprising the surfactant and other components of the composition of g. sodium carbonate and builder. Hitherto, the level of anionic surfactant present in such adjuncts has been limited due to the need to provide good flow properties and reduce the tendency to agglomerate.

EP-A-506 184 (Unilever) discloses a process for the continuous dry neutralisation of liquidacid precursor of anionic surfactant. Detergent particles having an active detergent content of 30 to 40% by weight may be prepared by this process.

EFP 572 957 discloses a process for producing a powdery anionic surfactant by feeding an aqueous slurry of the surfactant containing 60 to 50% solids into an evaporator, forming a film of the surfactant on the reactor wall and screpping if from the wall whilst drying and concentrating the slurry.

It is disclosed that the reactor wall is at a temperature of 50 to 140°C; 130°C is the highest wall temperature exemplified. Higher temperatures are said to cause thermal degradation and colour tone change and are thus disact-vantageous. Further, EP 572 957 discloses that the blades in the reactor are operated to provide at its seed of preferably 2 to 20 m/s, with 10.5 m/s being the highest tip speed which is exemplified. Bulk densities of up to about 0.5 g/cm² am disclosed.

We have found that contrary to the teaching of the prior at detegent particles having a high bulk density, a high level of anionic surfactant and excellent powder properties may be produced by heating a paste containing the surfactant in a first zone at a higher temperature than hitherto envisaged in the art and then cooling the thus formed particles. Moreover process throughput may be increased and the level of fine material and the particle size distribution of the particles may be carefully controlled.

Accordingly a first aspect of the invention provides a process for the production of detergent particles comprising at least 75%, prefereby at least 65% by weight of an anionic surfactant and no more than 10% by weight of water which comprises feeding a paste material comprising water in an amount of more than 10% by weight of the paste and the surfactant into a drying zone, healing the paste material to a temperature in excess of 130°C and preferably in access of 140°C in the said drying zone to reduce the water content to not more than 10% by weight and subsequently cooling the material in a cooling zone to form detergent particles wherein at least 80% of the particles have a particle size of 180 to 1500 µm and less than 10% have a particle size of 180 to 1500 µm and less than 10% have a particle size less than 180 µm.

Desirably the drying zone is under a slight vacuum to facilitate the removal of water and volatiles. The vacuum may be from 100 Torr up to atmospheric pressure as this provides significant process fate/billing. However, a vacuum in excess of 500 Torr up to atmospheric has the advantage of reducing capital investment whilst providing vacuum constitor.

We have found that Improved control of residence time and particle size may be secured and process throughput may be increased by adjusting the material in the drying and/or cooling zone.

50 Preferably, the paste is agitation with agitation means which have a tip speed in excess of 15 ms<sup>-1</sup> and preferably in excess of 20 ms<sup>-1</sup>

The process is preferably continuous as this facilitates continuous transportation of the particles. In a continuous process the flow rate is suitably of the order of 10 to 25 kg/m<sup>2</sup>/hr and preferably 17 to 22 kg/m<sup>2</sup>/hr e.g. 20 kg/m<sup>2</sup>/hr.

Suitably the average residence time in the drying zone is less than 5 minutes. A residence time of less than 4 minutes is especially preferred with as low a residence time as possible being most preferred.

Agitation of the paste in the healing zone generally provides efficient heat transfer within the paste and facilitate recorded of water. Agitation reduces the contact time between the paste particles and the wall of the drying zone which together with efficient heat transfer, reduces the likelihood of hot spots forming which may lead to decomposition.

Moreover, improved drying is secured thus allowing a shorter residence time/increased throughput in the drying zone.

To avoid thermal decomposition, the paste material is preferably not heated to a temperature in excess of 170°C.

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The process of the present invention permits the formation of particles having a high bulk density for example in excess of 150 g/cm<sup>3</sup>.

The material is cooled in a cooling zone which is suitably operated at a temperature not in excess of 50°C and preferably not in excess of 40°C e.g. 30°C. Desirably there is agitation within the cooling zone to provide efficient cooling of the material therein. By actively cooling the particles, the possibility of thermal decomposition occurring due to the particles being heated to a higher temperature than previously disclosed, is reduced and the tackiness of the particles may be reduced. Such active cooling may be through circulation of, for example, cold water or liquid nitrogen around the cooling zone, for example, in a cooling packet.

The paste material preferably comprises a mixture of anionic surfactant and water although other components may be present if desired or carried through as impurities from an up-stream process, for example production of the surfactant. Preferably the past enaberial comprises at least 50% by weight, more preferably at least 56% and ospecially at least 70% by weight of anionic surfactant. Suitably the paste comprises no more than 40% and preferably no more than 30% by weight of water. The paste material should be pumpable at the temperature at which it is to be fed into the driving zone and this may kimit the maximum they of surfactant process therein.

The paste is suitably fed to the drying zone at a temperature of 50 to 70°C and preferably 50 to 65°C where the paste comprises PAS, LES and/or LAS.

The process of the invention may be carried out in any suitable apparatus however it is preferred that a flash reactor is employed. Suitable flash reactors include e.g., the Flash Drier system available from VFNSpA processi impirant industriali. Desirably drying zone has a heat transfer area of at least 10m². The cooling zone desirably has a heat transfer area of at least at 5m².

Optionally two or more drying zones may be employed before the cooling zone as desired. A single apparatue may be employed to provide the drying zone and cooling zone as desired or alternatively separate apparatus for example a drier and a cooling fluid bed may be employed.

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Suitably the drying zone is substantially circular in cross section and is thus defined by a cylindrical vall. Preferably the said wall is heated by means of a heating jacket through which water, steam or oil may be feet. The inside of the said wall is preferably maintained at a temperature of at least 130°C and especially at least 140°C. Preferably the drying zone has an evaporation rate of 3 to 25, and especially 5 to 20 kg water per riff of heat surface per hour.

The cooling zone is preferably defined by a cylindrical wall. Where the process is continuous, the apparatus is suitably arranged such that the drying zone and cooling zone are substantially herizontally aligned to facilitate efficient drying, cooling and transport of the material through the drying and cooling zones in a generally horizontal direction.

Suitably the drying zone and preferably the cocling zone have aglistion means therein which aglistes and transports the surfactant paste and forming granules through the said zones. The aglistion means preferably comprises a series of radially extending pacidies and/or brades mounted on an axially mounted rotatable shaft. Desirably the pacidies and/or brades are inclined in order to effect transportation and preferably have a clearance from the inner wall of no more than 10mm, for example 5mm.

We have found that the present invention has especial applicability in the production of detergent particles comprising PAS. PAS is presently available on the market in tine powder form or in nocole form. The fine powder is generally dusty, having a significant quantity of particles of less than 150 micrometres. PAS nocoles are generally produced by axtruding dried PAS which has the appearance of soap chips and typically have a very large particle size and a very low porosity leading to poor dissolution characteristics. To increase the level of detergent active material in a detergent composition it is known to post-foxes detergent activates to provide a composition having a high level of active material.

However, PAS in fine powder form and PAS noodles are generally not suitable for post-dosing into a detergent composition as the composition particles and the post-dosed particles are generally of different particle size and thus tond to segregate and be unsightly. The process according to the present invention enables detergent particles having a high level of detergent active material and suitable prograft and enable size detargents to be obtained.

Accordingly a third aspect of the invention provides delargent particles comprising at least 75 % by weight of the particle of an anionic surfactant, preferably PAS, and not more than 10% by weight of the particle of water, the particles being obtainable by a process according to the first or second aspect of the invention.

According to a fourth aspect of the invention there is provided detergent particles comprising an anionic surfaciant, preferably PAS and preferably FAS and particle size of stribution such that at least 80% of the particles have a particle size of 180 to 1950 micrometres, preferably 280 to 1200 micrometres and less than 10% and preferably less than 5% of the particles have a particle size of 180 to 1950 micrometres.

Suitably the anionic surfactant in the detergent particles is present in an amount of at least 55% preferably at least 90% and desirably at least 94% by weight of the particles. It is desirable that the particles also comprise water in amount of 1 to 10% and preferably 1 to 8% by weight of the particles. The water in the particle provides improved

granule integrity thus reducing the level of the fine particles.

Suitably at least 80%, preferably 90% and more preferably 95% of the particles have a mean particle size of 300 to 1000 micrometres and more preferably 400 to 900 micrometres.

Desirably the detergent particles have an aspect ratio not in excess of 2 and more preferably are generally spherical in order to reduce aggregation from other particles in a formulated detergent composition and to enhance the visual appearance of the powder.

Suitably the PAS surfactant has a chain length of  $C_{10}$  to  $C_{22}$  preferably  $C_{12}$  to  $C_{18}$  and more preferably a narrow range of  $C_{12}$  to  $C_{14}$ . Coco PAS is particularly desirable

The detergent particle may comprise mixtures of PAS with other surfactants and/or non surfactant components as desired.

Suitable other surfactants may comprise alkyl benzene sulphonates, oxo alcohol sulphates for example  $C_{11}$  to  $C_{15}$  and  $C_{12}$  for  $C_{13}$  for  $C_{14}$  for  $C_{15}$  and  $C_{15}$  for  $C_{15}$ 

Especially preferred are PAS rich particles, that is particles in which the amount of PAS exceeds the amount of any other surfactant or non-surfactant and more preferably exceeds the total amount of all other surfactant components.

Generally the sodium salt of the surfactants will be employed.

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In general PAS active has poorer water-solubility characteristics than other commonly used detergent-cityes, for example LAS. This is especially the case all ower water temperatures which are typically employed for fatior watering in some countries. The dissolution characteristics for detergent particles comprising PAS and especially PAS-rich particles are influenced by such characteristics it is desired to improve the solubility thereof especially in cold waterior to reduce problems of deposition of the undissolved material on washed fabrics and wash inefficiency through non-participation of some carticles in the washing process.

We have found that the dissolution characteristics of particles comprising PAS may be improved by reducing the Krafft temperature of the PAS active to below 13°C, the Krafft temperature for PAS which is conventionally employed in deterent products.

Accordingly, a further aspect of the invention provides detergent particles comprising PAS, preferably at a level of at least 50%, more preferably at least 70% and especially at least 85% by weight of the particle, wherein the PAS has a Krafft temperature below 13°C and the average particle size is from 180 to 1500 µm.

Preferably the particles are produced by a process according to the invention as herein described.

Suitably at lease 50% and especially at least 70% of the PAS has a linear alkyl chain.

Preferably the Krafft temperature is below 10°C and more preferably below 5°C as the solubility of the PAS is significantly superior at temperatures above the Krafft temperature.

The Krafft temperature of the PAS may be reduced by any suitable means.

It has been surprisingly found that by employing a narrow alkly chain length distribution, the Krafft temperature thereof may be reduced. Preferably at least 90% and preferably at least 95% of the PAS active has a chain length of C<sub>2</sub>, to C<sub>16</sub>, and especially, for example EMPICOL LXV100 (tradename) av, Albright and Wilson

The Krafft temperature may also be reduced by employing a branched detergent active, preferably an alkylobenzene sulphonate, alcohord sulphate. Geurbot alcohord sulphate, secondary and preferably premixing together with a linear alcohol sulphate. Branched chain surfactants may assist in loam deen station which is deskribed to the consumer in some markets.

Examples of suitable branched surfactants include PETRELAB 550, LIAL 123 AS (ex DAC).

The Krafft temperature of the PAS may be reduced by employing a quaternary ammonium countarion to rup to 50 mole%, preferebly up to 50 mole% and aspecially up to 20 mole% of the determined nature in particles. Preferably the quaternary ammonium counterion is selected from ammonium and quaternised mono, di or tri alkanol amine, for example athenol amines.

Examples of suitable materials include the TEXAPON (tradename) range of surfactants ex Henkel,

The Krafft temperature may be lowered by employing, a narrow chain length distribution, a branched chain surlactant or a quaternary ammonium counterion, preferably a combination of these factors is employed to achieve further improvement in the solubility of the detergent particles

Other non-surfactant components which may be present in the detergent particles include dispersion acks, preferably polymeric dispersion acks and more preferably nos, sugars, polysigh/enexides, and builders as hereinfert described. If desired the detergent particles may comprise an organic and/or inorganic sail. Suitable materials in salts, preferably sodium, or throughoughouth, citrates, carbonates, subchaste, chlorides.

It is especially preferred that a salt be present in the particle when the anionic surfactant comprises LAS.

The salt may be present at a level of up to 40% and preferably up to 30% by weight of the particles.

The detergent particles may be post-dosed directly to a base powder obtained from any conventional detergent production process including a non tower process in which the components of the detergent composition are mixed and

granulated as described og in EP-A-587 339 (Unitever) and a spray drying process optionally followed by a post tower densification. As the detergent particles produced by the present invention may be post-dosed to such powders a significant degree of formulation flexibility is obtained and the level of active material in the fully formulated composition may be very high as desired. A further advantage is that a base powder which is substantially free of detergent active compounds may be produced as the detergent active compounds may be introduced substantially wholly as post-dosed particles.

Accordingly a further aspect of the invention provides a detergent composition comprising detergent particles according the third or fourth aspects of the invention and a base powder.

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The option of reducing the level of detergent active material in a base powder is especially advantageous where the base powder is produced by a spray drying process as a lower level of detergent active compound in the spray drying process permits a higher throughout to be secured thus increasing overall production efficiency.

Compositions according to the fifth aspect of the invention generally contain, in addition to the detergent-active compound, a detergency builder and optionally bleaching components and other active ingredients to enhance performance and proporties.

Detergent compositions of the invention may contain, in addition to the post-dosed detergent particles, one or more detergent-active compounds (surfactants) which may be chosen from soep and non-scap anionic, calloricia, no-noincia, amphotrier and zwitterionic detergent-active compounds, and mixtures thereof. Many suitable detergent-active compounds are available and are fully described in the literature, for example, in "Surface-Active Agents and Detergents", Volumes i and II, by Schwartz. Perry and Bench. The preferred detergent-active compounds that can be used are soaps and synthetic non-soap anions and nonionic compounds.

Anionic surfactants are well-known to those skilled in the art. Examples include alkylbenzene sulphonates, particularly linear alkylbenzene sulphonates having an alkyl chain length of C8-C15, primary and secondary alkyl sulphates, particularly C12-C15 primary alkyl sulphates, alkyl ether sulphates, olefin sulphonates; alkyl xylene sulphonates, dialkyl sulphosuconates; and latty acid ester sulphonates. Sodium sails are generally preferred

Nonionic surfactants that may be used include the primary and secondary alcohol ethoxylates, especially the C<sub>8</sub>-C<sub>20</sub> allphatic alcohols ethoxylated with an average of from 1 to 20 moles of ethylene oxide per mole of alcohol, and more especially the C<sub>10</sub>-C<sub>12</sub> primary and secondary allphatic alcohols ethoxylated with an average of from 1 to 10 moles of athylene oxide per mole of alcohol. Non-ethoxylated nonionic surfactants include alkylpolyglycosides, glycerol monoeithers, and oxphivtydroxymaties follocamics.

The total amount of surfactant present in the detergent composition is suitably from 5 to 40 wt% aithough amounts outside this range may be employed as desired.

The detergent compositions of the invention generally also contain a detergency builder. The total amount of detergency builder in the compositions is suitably from 10 to 80 w/s, preferably from 15 to 60 w/s. The builder may be present in an adjunct with other components or, if desired, separate builder particles containing one or more builder materials may be employed.

Inorganic builders that may be present include sodium carbonate, if desired in combination with a crystallisation seed for calcium carbonate, as disclosed in GB 1 473 950 (Unilever); crystalline and amorphous aluminos licates, for example, zeolities as disclosed in GB 1 473 201 (Henkel), amorphous aluminosilicates as disclosed in GB 1 473 202 (Henkel) and mixed crystalline/amorphous aluminosilicates as disclosed in GB 1 470 205 (Procter & Gamble); and layered silicates as disclosed in GB 1 470 205 (Procter & Gamble); and layered silicates as disclosed fixed processing the companies of the control of

Zoolte builders may suitably be present in an amount of from 10 to 60 w% and preferably an amount of from 15 to 50 w%. The zeelfe used in most commercial particulate detergent compositions is zeolite A. Advantageously, however, maximum aluminium zeolite P (zeolite MAP) described and claimed in EP 984 0704 (Unilever) may be used. Zeolite MAP is an alkali metal aluminosilicate of the P type having a silicon to aluminium ratio not exceeding 1.33, preferably not exceeding 1.50, and more preferably not exceeding 1.50.

Organic buildors that may be present include polycarboxylete polymers such as polyacrylates, acrylicimaleic ocpolymers, and acrylic phosphinates, monomeric polycarboxyletes such as citrates, gluconates, oxydieuccinates, glycerol mono-, di-and trisuccinates, carboxymethyloxysuccinates, earboxymethyloxymalonates, dipioiniates, brydroxyethyliminodiacetates, alkyl- and alkenylmalonates and succinates; and sulphonated fatty acid saits. A copolymer of maloic acid, acrylic acid and vinyl acetate is especially preferred as it is biodegradable and thus environmentally destrable. This list is not intended to be exhaustive.

Especially preferred organic builders are cirates, suitably used in amounts of from 5 to 30 wt%, preferably from 10 to 25 wt%, and acrylic polymers, more especially acrylic/maleic copolymers, suitably used in amounts of from 0.5 to 15 wt%, preferably from 1 to 10 wt%. The builder is preferably present in alkali metal salt, especially socium salt, for

Suitably the builder system comprises a crystalline layered silicate, for example, SKS-6 ex Hoechst, a zeolite, for example, zeolite A and optionally an alkali metal citrate.

Detergent compositions according to the invention may also contain a bleach system, desirably a peroxy bleach

### FP 0 777 719 R1

compound for example, an inorganic perselt or organic peroxyscid. capable of yielding hydrogen peroxicle in aqueous solution. The peroxy bleach compound may be used in conjunction with a bleach activator (bleach precursor) to improve bleaching action at low wesh temperatures. An especially preferred bleach system comprises a peroxy bleach compound (preferably sodium percarbonate optionally together with a bleach activator), and a transition metal bleach catavit as described and oblamed in EP 458 3974. EP 458 3984 and EP 509 7974 (Unilever).

The compositions of the invention may contain alkall metal, preferably sodium, carbonate, in order to increase detergency and ease processing. Sodium carbonate may suitably be present in an amount from 1 to 60 Mt%, preferably from 2 to 40 Mt%. However, compositions containing filtle or no sodium carbonate are also within the scope of the invention.

Powder flow may be improved by the incorporation of a small amount of a powder structurant, for example, a fatty add soap), a sugar, an acrylate or acrylate/maleate polymer, or sodium silicate which is suitably present in an amount of from 1 to 5 wt%.

Other materials that may be present in detergent compositions of the invention include sodium silicate, antiredeposition agents such as cellulosis polymers, fluorescers, inorganic salts such as sodium sulphate, lather control agreor lather boosters as appropriate, proteolytic and lipolytic enzymes; dyes: coloured speckles; perfumes; foam controllers; and fabric softening compounds. This list is not intended to be exhaustlyen.

The base composition is suitably prepared by spray-drying a sturry of compatible heat-insensitive ingredients, and then spraying on, admixing and/or postdosing those ingredients unsuitable for processing via the sturry. The detergent particles produced according to the process of the present invention are post-dosed to the base composition by conventional methods.

Detergent compositions of the invention preferably have a bulk density of at least 500 g/l, more preferably at least 550 g/litre, more preferably at least 700 g/litre.

Such powders may be prepared either by spray-drying, by post-tower densification of spray-dried powder, or by wholly non-tower methods such as dry mixing and gramulation. A high-speed mixer/gramulator may advantageously be used for such mixing. Processes using high-speed mixer/gramulators are disclosed, for example, in EP 340 013A. EP 340 3151 Au file 140 01374. Unlieven.

The invention is illustrated by the following non-limiting Examples.

### Example 1

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An aqueous paste comprising 70% by weight of sodium cocoPAS was fed into the drying zone of a Flash Drier manufactured by VRVSpA. Italy at a temperature of 60°C. A small vactuum was applied to the drying zone The initial throughput in the Flash Drier was 120 kg/hr of paste. The temperature of the wall of the drying zone was initially 140°C. The heat trensfer area of the drying and cooling zones was respectively 10mg and 5mg.

The temperature of the wall of the drying zone was raised in steps to 170°C. Correspondingly the throughput was increased to 430 kg/hr at 170°C. At each step the process conditions were stabilised for 15 minutes. The particles then passed to a cooling zone operated at a temperature of 30°C.

PAS granules comprising at least 85% PAS and 51 o 8.5% moisture content were obtained from each step having a bulk density of at least 550 glf with a dynamic flow rate of at least 100 m/ls. The samples obtained all had a whiteness of at least 80 on a gray-scale of 1 (black) to 100 (white) measured using a Minotla CR-210 Chromamoter. Commercially available detergent powders generally have a whiteness of at least 75 thus illustrating that little thermal decomposition leading to policy impatiment had occurred.

The samples all contained less than 5% fine material (180  $\mu m$  or less) and the particles exhibited good strength and low friability.

The solubility of the samples was tested in water at 20°C, conductivity being employed to measure the % dissolution. In all cases, at least 75% of the sample had dissolved after 20 seconds and at least 95% after 30 seconds illustrating the excellent solubility characteristics of the PAS granules.

#### Example 2

The procedure of Example 1 was repeated using a small scale Flash Orier having a heat transfer surface area of 0.5m<sup>2</sup> (dying area: cooling area = 2.1). The aglitator (ip speed was about 90m<sup>2</sup>, and the drying and cooling zones were operated at temperatures of 160°C and 40°C respectively. PAS granules containing 95 to 97.5% PAS and 1.5 to 2% moisture were obtained.

## 55 Example 3

An example of a detergent composition according to the invention is listed below in which the base powder, PAS granules and other components are dry-mixed:

Base Powder		60%
- Nonionic surfactant	12	
- Soap	2	
- Zeolite builder	38	
- Moisture, salts, NDOM	8	
PAS granules		9%
Percarbonate		20%
Minors (include foam suppressor, TAED, enzyme)		11%

The composition exhibited good detergency and dissolution characteristics

### Example 4

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A series of compositions containing LAS as the detergent actives were produced as detailed in the following Table, by the procedure outlined below.

	Product Composition as Weight %					
LAS	75	77.0	79 5	75 6	68.2	
Water	2.0	2.0	2.0	2.0	2.0	
Sodium Citrate	0.0	0.0	8.9	18.9	0.0	
STP	20	17.6	6.3	0.0	26.8	
NDOM and minors	3.0	3.4	3.3	3.5	3.0	

An aqueous paste of the components of the compositions was fed into the drying zone of a flash-drier having a heat transfer surface of 1.2m² (drying area:cooling area 2:1).

The agitated tip speed was about a 50ms 1 and the drying and cooling zones were operated at temperatures of about 160°C and 20°C respectively. The pastes were fed into the dryer at a feed rate of 10 to 30 kghr1.

Solid particles containing LAS and a salt were produced satisfactorily.

## Example 5 to 11

Examples of detergent particles according to the present invention and which may be produced by a process according to the present invention are listed in the following Table.

The time for 90% of the particles to dissolve in water at 5°C was measured using an AGB-4001 conductivity meter with a final surfactant concentration of 0.2 ct<sup>-1</sup> in demineralised water.

	5	6	7	8	9	10	11
Coco PAS C <sub>12-14</sub> sodium salt	100	50	80	70	BO	90	76
aTEA salt	-	-	-	10	20	10	-
blial 123 AS	-	50	20	20	-	-	19
°PEG 4000	-	-	-		-	-	5
Dissolution time (mins)	20	1	3	1	0.5	1	3

<sup>b</sup> branched PAS sodium salt ex DAC

c ex BDH

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#### FP 0 777 719 R1

#### Claims

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- A process for the production of detergent particles comprising at least 75% by weight of an anionic surfactant and
  no more than 10% by weight of water which comprises feeding a paste material comprising water in an amount
  of more than 10% by weight of the paste and the surfactant into a divigor zone, heating the paste melerial to a
  temperature in excess of 130°C in the said drying zone to reduce the water content to not more than 10% by weight
  and subsequently actively actively acolling the material in a cooling zone to form detergent particles wherein at least 80°C.

  of the particles have a particle size of 150°D 150°O um and less than 10% have a particle size less than 180°C.
- The process of claim 1, further comprising agitating the paste with agitation means which have a tip speed in excess of 15 ms<sup>-1</sup>.
  - 3. A process according to claim 1 or 2 in which the anionic surfactant comprises primary alkyl sulphate.
- A process according to any preceding claim wherein the anionic surfactant is present in the particles at a level of at least 90% by weight of the particles.
  - A process according to any preceding claim wherein the paste material is heated to a temperature in excess of 140°C
  - Detergent particles comprising at least 75 % by weight of the particle of an anionic surfactant, and not more than 10% by weight of the particle of water, the particles being obtainable by the process according to claim 1.
- Detergent particles obtainable by the process of claim 1 and comprising PAS wherein the PAS has a Krafft temperature below 13°C and preferably below 10°C.
  - 8. Detergent particles according to claim 7 wherein the PAS has an alkyl chain length of C12 to C18.
  - 9. Detergent particles according to claim 7 or 8 which further comprise a branched chain anionic surfactant.
  - Detergent particles according to claims 7 to 9 wherein the counterion of up to 50 mole% of the PAS is a quaternary ammonium ion.
  - Detergent particles according to any one of claims 5 to 10 wherein the particles have a porosity of 5 to 50% volume
    of the particle.
  - A detergent composition comprising detergent particles as defined in any one of ctaims 6 to 11 and a base powder comprising a surfactant and/or a builder and, optionally, further comprises an alkali metal percarbonate.
- 40 13. A detergent composition according to claim 12 in which the base powder comprises a builder comprising a silicate, preferably a crystalline layered silicate, and/or a zeolite.

# Patentansprüche

- 1. Verfahren zur Herstellung von Waschmitteltellichen, umfassend mindestens 75 Gew.-% eines anleinischen Tensids und nicht mehr als 10 Gew.-% Wassen umfassend Zufuhr eines Pasten-Haberials, das Wasser in einer Mange von mehr als 10 Gew.-% der Paste und das Tensids enthält, in eine Trockenzone, Erhitzen das Pasten-Materials auf eine Temperatur oberhalb 13 DrC in der Trockenzone, um den Wasseranteil auf hicht mehr als 10 Gew.-% zu verminden und anschließend aktikt Wölhel des Materials in einer Kühlzone zur Herstellung von Waschmittelleichen, worin mindestens 80% der Teilchen eine Teilchengröße von 180 bis 1500 µm aufweisen und weniger als 10% eine Teilchengröße von weniger als 150 µm aufweisen.
- Verlahren nach Anspruch 1, das außerdem Bewegen der Paste mit einer Bewegungsvorrichtung, die eine Spitzengeschwindigkeit von mehr als 15 m·s<sup>-1</sup> aufweist, umfaßt.
  - 3. Verfahren nach Anspruch 1 oder 2, wobei das anionische Tensid primäres Alkylsulfat umfaßt.

- Verfahren nach einem vorangehenden Anspruch, wobei das anionische Tensid in den Teilchen in einer Menge von mindestens 90 Gew.-% der Teilchen vorliegt.
- Verfahren nach einem vorangehenden Anspruch, wobei das Pasten-Material auf eine Temperatur oberhalb 140°C erhitzt wird.
  - Waschmittelteilichen, wobei mindestens 75 Gew.-% der Teilchen ein anionisches Tensid und nicht mehr als 10 Gew.-% der Teilchen Wasser umfassen, und die Teilchen durch das Verfahren nach Anspruch 1 erhältlich sind
- Waschmittelteilchen, erhältlich durch das Verlahren nach Anspruch 1 und umfassend PAS, wobei das PAS eine Krafit-Temperatur unterhalb 13°C und vorzugsweise unterhalb 10°C aufweist.
  - 8. Waschmittelteilchen nach Anspruch 7, wobei das PAS eine Alkylkettenlänge von C12 bis C16 aufweist
- 9. Waschmittelteilchen nach Anspruch 7 oder 8, die außerdem ein verzweigtkettiges anionisches Tensid umfassen.
  - Waschmittelteilchen nach Ansprüchen 7 bis 9, wobei das Gegenion von bis zu 50 Mol-% des PAS ein quatemäres Ammoniumien ist.
- Waschmittettelichen nach einem der Ansprüche 5 bis 10, wobei die Teilchen eine Porosität von 5 bis 50 Volumen-% der Teilchen aufweisen
  - Waschmittel, umfassend Waschmittelteilchen nach einem der Ansprüche 6 bis 11 und ein Grundpulver, das ein Tensid und/oder einen Builder umfaßt und gegebenenfalls außerdem ein Alkalmetallipercarbonat umfaßt
    - Waschmittel nach Anspruch 12, wobei das Grundpulver einen Builder, umfassend ein Silicat, vorzugsweise ein kristallines Schichtsilicat und/oder einen Zeolith, umfaßt.

# 30 Revendications

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- Un procédé de production de particules clétergentes comprenant au mon » 75 % en masse d'un agent tensionabil ainoinque et au plus 10 % en masse d'au comprand le chargement dans la zone de séchage d'un matériau on pâte comprenant de l'eau dans une quantité ne dépassant pas 10 % en masse de la pâte et de l'agent tensioactif, de chauffage du matériau en pâte à une température aupérieure à 130 °C dans la citiez cone de séchage afin de réduire le contenu en eur au puls 10 % en masse et par la suite le refrocissement actif du matérau dans une zone de réfroitssement afin de former des particules détergentes dans lesquelles au moins 80 % des particules présentent une taile de particule comprise entre 180 et 1500 μm, et moins de 10% présentent une taille de particule inférieure à 180 μm.
  - Le procédé de la Revendication 1 comprenant en plus une agitation de la pête à l'aide d'un moyen d'agitation avec une vitesse périphérique de l'hélico supérieure à 15 ms<sup>-1</sup>.
- Un procédé en accord avec la Revendication 1 ou 2 dans laquelle l'agent tensioactif anionique comprend un sulfate d'alkyle primaire.
  - Un procédé en accord avec l'une quelconque des revendications précédentes dans laquelle l'agent tensioactif anionique est présent dans les particules à un niveau au moins égal à 90 % en masse des particules.
- Un procédé en accord avec l'une quelconque des revendications précédentes dans laquelle le matériau en pâte est chauffé à une température subérieure à 140 °C.
  - Des particules détergentes comprenant au moins 75 % en masse des particules d'un agent tensicactif anionique, et au plus 10 % en masse de la particule d'eau, les particules pouvant être obtenues selon le procédé en accord avec la Rovendication.
    - 7. Des particules détergentes pouvant être obtenues selon le procédé de la Revendication 1 et comprenant du PAS, cans laquelle le PAS a une température de Krafft inférieure à 13 °C et de préférence inférieure à 10 °C.

- Des particules détergentes en accord avec la Revendication 7 dans laquelle le PAS a une longueur de chaîne alkyle comprise entre C<sub>12</sub> et C<sub>16</sub>.
- Des particules détergentes en accord avec la Revendication 7 ou 8 qui comprend en plus un agent tensicactif anionique de chaîne ramifiée

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- 10. Des particules détergentes en accord avec les Revendications 7 à 9 dans fesquelles le contre-ion comprenant jusqu'à 50 % en mole du PAS est un ion d'ammonium quaternaire.
- 11. Des particules détergentes en accord avec l'une quelconque des Revendications 5 à 10 dans lesquelles les particules ont une corosité comprise entre 5 et 50 % du volume de la particule.
  - 12. Une composition détergente comprenant des particules détergentes telles que définies dans l'une quelconque des Revendicators 6 à 11 et une poudre de base comprenant un agent tensioactif étique un édificateur, et en option comprenant un percabonate de métal alcalin
    - 13. Une composition détergente en accord avec la Revendication 12 dans laquelle la poudre de base comprend un édificateur comprenant un silicate, de préférence, un silicate stratifié cristallin, et/ou une zéolite.